09/941,882

(230,786.1)

FILE 'HOME' ENTERED AT 15:52:35 ON 15 JUL 2003

=> file biosis medline caplus wpids uspatfull

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FILE 'WPIDS' ENTERED AT 15:52:59 ON 15 JUL 2003 COPYRIGHT (C) 2003 THOMSON DERWENT

FILE 'USPATFULL' ENTERED AT 15:52:59 ON 15 JUL 2003 CA INDEXING COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

*** YOU HAVE NEW MAIL ***

=> s nucleic acid? sequencing

L1 2960 NUCLEIC ACID? SEQUENCING

=> s l1 and heat

L2 1048 L1 AND HEAT

=> s 12 and diphenyliodonium

L3 1 L2 AND DIPHENYLIODONIUM

=> d 13 bib abs

L3 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS

AN 2002:736776 CAPLUS

DN 137:258477

TI Real-time nucleic acid sequencing by monitoring of labeled base incorporation followed by removal and substitution of the labeled nucleotide

IN Williams, Peter; Taylor, Thomas J.; Williams, Daniel J. B.; Gould, Ian; Hayes, Mark A.

PA USA

SO U.S. Pat. Appl. Publ., 37 pp., Cont.-in-part of U.S. Ser. No. 673,544. CODEN: USXXCO

DT Patent

LA English

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	US 2002137062	A1	20020926	US 2001-941882	20010828
	WO 9957321	A1	19991111	WO 1999-US9616	19990430
	W: CA, JP,	US			

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

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WO 2003020895
                            20030313
                                           WO 2002-US27605 20020828
                      A2
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
             CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
             NE, SN, TD, TG
PRAI US 1998-83840P
                       Ρ
                            19980501
     WO 1999-US9616
                       Α2
                            19990430
     US 2001-673544
                       A2
                            20010226
     US 2001-941882
                       Α
                            20010828
AΒ
     The present invention relates to a novel method for analyzing nucleic acid
     sequences based on real-time detection of DNA polymerase-catalyzed
     incorporation of each of the four nucleotide bases, supplied individually
     and serially in a microfluidic system, to a reaction cell contg. a
     template system comprising a DNA fragment of unknown sequence and an
     oligonucleotide primer. Incorporation of a nucleotide base into the
     template system can be detected by any of a variety of methods including
     but not limited to fluorescence and chemiluminescence detection.
     Alternatively, microcalorimetric detection of the heat generated
     by the incorporation of a nucleotide into the extending template system
     using thermopile, thermistor and refractive index measurements can be used
     to detect extension reactions. The preferred method uses a two-stage
     primer extension procedure. In the first stage the primer is extended by
     exposing it to labeled derivs. of nucleotide triphosphates in the presence
     of an exonuclease free DNA polymerase until a signal indicating
     incorporation is seen. THe labeled nucleotide is removed and replaced
     using the correctly identified nucleotide and a mixt. of DNA polymerases
     with and without exonuclease activity. The use of microcalorimetry to
     detect base incorporation is demonstrated. The use of
     diphenyliodonium to bleach fluorescent labels is demonstrated.
=> d his
     (FILE 'HOME' ENTERED AT 15:52:35 ON 15 JUL 2003)
     FILE 'BIOSIS, MEDLINE, CAPLUS, WPIDS, USPATFULL' ENTERED AT 15:52:59 ON
     15 JUL 2003
           2960 S NUCLEIC ACID? SEQUENCING
L1
           1048 S L1 AND HEAT
L_2
L3
              1 S L2 AND DIPHENYLIODONIUM
=> s 12 and pyrophosphate
           228 L2 AND PYROPHOSPHATE
L4
=> s 14 and hydrolosis (4a) pyrophosphate
             0 L4 AND HYDROLOSIS (4A) PYROPHOSPHATE
=> s 14 and hydroly? (4a) pyrophosphate
L6
             3 L4 AND HYDROLY? (4A) PYROPHOSPHATE
=> d 16 bib abs 1-3
1.6
     ANSWER 1 OF 3 USPATFULL
AN
       2003:86313 USPATFULL
ΤТ
      Novel human 39228, 21956, 25856, 22244, 8701, 32263, 50250, 55158,
       47765, 62088, 50566, and 48118 molecules and uses therefor
```

```
Meyers, Rachel E., Newton, MA, UNITED STATES
IN
       Rudolph-Owen, Laura A., Jamaica Plain, MA, UNITED STATES
       Kapeller-Libermann, Rosana, Chestnut Hill, MA, UNITED STATES
       Millennium Pharmaceuticals, Inc., Cambridge, MA, UNITED STATES, 02139
PA
       (U.S. corporation)
       US 2003059919
                               20030327
PΙ
                          A1
       US 2002-160501
                               20020530 (10)
AΙ
                          A1
       Continuation-in-part of Ser. No. US 2001-838573, filed on 18 Apr 2001,
RLI
       PENDING Continuation-in-part of Ser. No. US 2001-870133, filed on 29 May
       2001, PENDING Continuation-in-part of Ser. No. US 2001-870130, filed on
       29 May 2001, PENDING Continuation-in-part of Ser. No. US 2001-862535,
       filed on 21 May 2001, PENDING Continuation-in-part of Ser. No. US
       2001-870383, filed on 29 May 2001, PENDING Continuation-in-part of Ser.
       No. US 2001-860821, filed on 18 May 2001, PENDING Continuation-in-part
       of Ser. No. US 2001-870110, filed on 29 May 2001, PENDING
       Continuation-in-part of Ser. No. US 2001-907509, filed on 16 Jul 2001,
       PENDING Continuation-in-part of Ser. No. US 2001-945327, filed on 31 Aug
       2001, PENDING
       US 2000-197747P
                           20000418 (60)
PRAI
                           20000526 (60)
       US 2000-207649P
       US 2000-207640P
                           20000526 (60)
       US 2000-205961P
                           20000519 (60)
       US 2000-207506P
                           20000526 (60)
       US 2000-205449P
                           20000519 (60)
       US 2000-207650P
                           20000526 (60)
       US 2000-218385P
                           20000714 (60)
       US 2000-229425P
                           20000831 (60)
       US 2001-318581P
                           20010910 (60)
\mathbf{DT}
       Utility
       APPLICATION
FS
       LAHIVE & COCKFIELD, 28 STATE STREET, BOSTON, MA, 02109
LREP
       Number of Claims: 23
CLMN
ECL
       Exemplary Claim: 1
DRWN
       100 Drawing Page(s)
LN.CNT 44311
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The invention provides isolated nucleic acids molecules, designated
AB
       39228, 21956, 25856, 22244, 8701, 32263, 50250, 55158, 47765, 62088,
       50566, and 48118 nucleic acid molecules, which encode novel GTPase
       activating molecules, cadherin molecules, and ankyrin containing family
       members. The invention also provides antisense nucleic acid molecules,
       recombinant expression vectors containing 39228, 21956, 25856, 22244,
       8701, 32263, 50250, 55158, 47765, 62088, 50566, and 48118 nucleic acid
       molecules, host cells into which the expression vectors have been
       introduced, and non-human transgenic animals in which a 39228, 21956,
       25856, 22244, 8701, 32263, 50250, 55158, 47765, 62088, 50566, or 48118
       gene has been introduced or disrupted. The invention still further
       provides isolated 39228, 21956, 25856, 22244, 8701, 32263, 50250, 55158,
       47765, 62088, 50566, and 48118 polypeptides, fusion polypeptides,
       antigenic peptides and anti-39228, 21956, 25856, 22244, 8701, 32263,
       50250, 55158, 47765, 62088, 50566, and 48118 antibodies. Diagnostic and
       therapeutic methods utilizing compositions of the invention are also
       provided.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
```

ANSWER 2 OF 3 USPATFULL

L6

```
2001:67439 USPATFULL
ΑN
       Mutant chimeric DNA polymerase
ΤI
       Gelfand, David Harrow, Oakland, CA, United States
IN
       Reichert, Fred Lawrence, Oakland, CA, United States
PA
       Roche Molecular Systems, Pleasanton, CA, United States (U.S.
```

```
corporation)
ΡI
       US 6228628
                          B1
                               20010508
       US 1998-105697
                               19980626 (9)
ΑI
PRAI
       US 1997-52065P
                           19970709 (60)
       Utility
DТ
FS
       Granted
EXNAM Primary Examiner: Achutamurthy, Ponnathapu; Assistant Examiner: Hutson,
       Richard
LREP
       Petry, Douglas A.
       Number of Claims: 63
CLMN
       Exemplary Claim: 1
ECL
       3 Drawing Figure(s); 8 Drawing Page(s)
DRWN
LN.CNT 2195
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AB
       Mutant, chimeric thermostable DNA polymerases are provided, along with
       purified DNA sequences that encode the enzymes. Also provided are
       methods for producing and using the enzymes.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 3 OF 3 USPATFULL
L6
AN
       97:81103 USPATFULL
TI
       Purified nucleic acid encoding a thermostable pyrophosphatase
IN
       Gelfand, David Harrow, Oakland, CA, United States
       Wang, Alice Ming, Lafayette, CA, United States
       Roche Molecular Systems, Inc., Branchburg, NJ, United States (U.S.
PΑ
       corporation)
       US 5665551
                               19970909
PΙ
       US 1995-528384
                               19950913 (8)
ΑI
       Utility
DT
FS
       Granted
EXNAM Primary Examiner: Hendricks, Keith D.
LREP
       Johnston, George W., Sias, Stacey R., Petry, Douglas A.
CLMN
       Number of Claims: 7
ECL
       Exemplary Claim: 7
DRWN
       No Drawings
LN.CNT 921
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Purified DNA sequences that encode a thermostable pyrophosphatase are
       provided. The purified DNA is obtained using a DNA probe consisting of
       SEQ ID NO: 1. Also provided are methods for producing thermostable
       pyrophosphatase.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
=> d 16 2-3 kwic
L6
     ANSWER 2 OF 3 USPATFULL
SUMM
       . . polymerase, methods for its synthesis, and methods for its use.
       The enzyme is useful in many recombinant DNA techniques, especially
       nucleic acid sequencing and nucleic acid
       amplification by the polymerase chain reaction (PCR).
SUMM
       Typically, sequencing by the chain termination method is carried out
       using repeated steps of primer extension followed by heat
       denaturation of the primer extension product-template duplex. This
       embodiment, referred to as cycle sequencing, is carried out in a
       thermal.
DETD
       The term "thermostable enzyme", as used herein, refers to an enzyme
       which is stable to heat and has an elevated temperature
       reaction optimum. The thermostable enzyme of the present invention
```

catalyzes primer extension optimally at a.

- DETD . . . catalyze the template-dependent incorporation of a deoxynucleotide onto the 3'-hydroxyl terminus of a primer, with the concomitant release of inorganic pyrophosphate (PPi). This polymerization reaction is reversible. DNA polymerases also catalyze the reverse reaction, pyrophosphorolysis, which is the degradation of DNA.
- DETD Inorganic pyrophosphatase (PPase), also known as pyrophosphate phosphohydrolase, catalyzes hydrolysis of inorganic pyrophosphate (PPi) to two molecules of orthophosphate. PPase plays an vital role in RNA and DNA synthesis in vivo. By cleaving.
- DETD . . . such as those derived from polyoma, adenovirus 2, bovine papilloma virus (BPV), or avian sarcoma viruses, or immunoglobulin promoters and **heat** shock promoters. A system for expressing DNA in mammalian systems using a BPV vector system is disclosed in U.S.
- DETD . . . polymerase is carried out in E. coli, which is a mesophilic bacterial host cell. Because E. coli host proteins are heat -sensitive, the recombinant thermostable DNA polymerase can be substantially enriched by heat inactivating the crude lysate. This step is done in the presence of a sufficient amount of salt (typically 0.2-0.4 M. . .
- DETD C. The amplified products from steps A and B are combined, heat denatured at 95.degree. C., annealed, and extended with DNA polymerase using standard techniques.
- DETD C. The amplified products from steps A and B were combined, heat denatured at 95.degree. C., annealed, and extended with DNA polymerase using standard techniques.
- DETD Ammonium sulfate was added to the Fraction II supernatant to a concentration of 0.4 M. Fraction II then was heat-treated as follows.
- DETD The heat treatment was carried out in a 3 liter Braun fermentor. The agitation rate was 250 rpm. The temperature was increased. . . over 6 minutes, held for 15 minutes, then cooled in the fermentor to 30.degree. C. as rapidly as possible. The heat -treated Fraction II supernatant from the PEI precipitation was removed from the fermentor and held on ice for at least 30. . .
- DETD . . . as a carrier. The DNA was precipitated by the addition of 1 ml 20% trichloroacetic acid (w/v) and 2% sodium pyrophosphate, and incubated at 0.degree. C. for 15 minutes. Precipitated DNA was collected on GF/C filter discs (Whatman International Ltd., Maidstone, England) and washed extensively with 5% trichloroacetic acid and 2% sodium pyrophosphate, then with 5% trichloroacetic acid, then with 5 ml of 95% ethanol, dried, and counted.

L6 ANSWER 3 OF 3 USPATFULL

- SUMM . . . relates to the in vitro synthesis of a thermostable pyrophosphatase. Thermostable pyrophosphatases are useful in many recombinant DNA techniques, especially nucleic acid sequencing and nucleic acid amplification by the polymerase chain reaction (PCR).
- SUMM Pyrophosphate is a common product of biosynthetic reactions.

 Inorganic pyrophosphatase (PPase), also known as pyrophosphate
 phosphohydrolase, catalyzes hydrolysis of inorganic
 pyrophosphate (PPi) to two molecules of orthophosphate. PPase
 plays an vital role in RNA and DNA synthesis in vivo. By cleaving.
- SUMM . . . between 25 and 40 times. Initial amplification conditions are chosen which favor the forward (polymerization) reaction (high dNTP concentrations, low pyrophosphate concentration). However, the amplification reaction results in an accumulation of pyrophosphate which increases the rate of the reverse reaction (pyrophosphorolysis), thereby decreasing the overall efficiency of the amplification reaction.

09567863

- SUMM The term "thermostable enzyme", as used herein, refers to an enzyme which is stable to **heat** and has an elevated temperature reaction optimum. The thermostable pyrophosphatase encoded by the nucleic acid sequence of the present invention. . .
- SUMM . . . such as those derived from polyoma, adenovirus 2, bovine papilloma virus (BPV), or avian sarcoma viruses, or immunoglobulin promoters and heat shock promoters. A system for expressing DNA in mammalian systems using a BPV vector system is disclosed in U.S.
- SUMM . . . PPase is carried out in E. coli, which is a mesophilic bacterial host cell. Because E. coli host proteins are heat -sensitive, the recombinant thermostable PPase can be substantially enriched by heat inactivating the crude lysate. This step is done in the presence of a sufficient amount of salt (typically 0.3M ammonium. . .
- SUMM Activity of the purified PPase is assayed as follows. The purified enzyme is used in the hydrolysis of inorganic pyrophosphate (PPi) with subsequent release of two orthophosphates, and resulting amount of orthophosphate is measured. The hydrolysis reaction is carried out. . .
- DETD . . . duty cycle for 3 minutes to reduce the viscosity. The sonicate was heated at 85.degree. C. for 15 minutes. The heat-treated supernatant was chilled rapidly to 0.degree. C., and the E. coli cell membranes and denatured proteins were removed following centrifugation.

=> d his (FILE 'HOME' ENTERED AT 15:52:35 ON 15 JUL 2003) FILE 'BIOSIS, MEDLINE, CAPLUS, WPIDS, USPATFULL' ENTERED AT 15:52:59 ON 15 JUL 2003 2960 S NUCLEIC ACID? SEQUENCING L1L21048 S L1 AND HEAT 1 S L2 AND DIPHENYLIODONIUM L3 228 S L2 AND PYROPHOSPHATE L4O S L4 AND HYDROLOSIS (4A) PYROPHOSPHATE L5 3 S L4 AND HYDROLY? (4A) PYROPHOSPHATE L6 => s l1 and destroy? (5a) fluorescen? (6a) remov? 0 L1 AND DESTROY? (5A) FLUORESCEN? (6A) REMOV? => s l1 and destroy? (6a) fluoresce? 1 L1 AND DESTROY? (6A) FLUORESCE? => d 18 bib abs ANSWER 1 OF 1 USPATFULL L894:100500 USPATFULL AN Method and apparatus for automatic nucleic acid sequence determination ΤI IN Tibbetts, Clark, Nashville, TN, United States Bowling, John M., Murfreesboro, TN, United States Vanderbilt University, Nashville, TN, United States (U.S. corporation) PΑ PΙ US 5365455 19941115 ΑI US 1991-763457 19910920 (7) DTUtility FS Granted EXNAM Primary Examiner: Harvey, Jack B.; Assistant Examiner: Choi, Jae H. Needle & Rosenberg LREP CLMN Number of Claims: 38 Exemplary Claim: 1 ECL DRWN 16 Drawing Figure(s); 16 Drawing Page(s) LN.CNT 1440 CAS INDEXING IS AVAILABLE FOR THIS PATENT. A method and system for automated nucleic acid sequence determination of AB a polynucleotide, wherein a nucleic acid sequencing ladder comprises signals corresponding to oligonucleotides formed from the polynucleotide, comprising the step of correlating, particularly in a trained neural network or a scatter plot, an intensity variable for each signal in the nucleic acid sequencing ladder with an informative variable for that signal, wherein the informative variable comprises information from at least two adjacent signals in the nucleic acid sequencing ladder, such that each signal in the nucleic acid sequencing ladder identified so as to determine the nucleic acid sequence corresponding to the polynucleotide. In particular, the relative separation between consecutive signals, the relative intensities between consecutive signals, and a pattern recognition factor, which incorporates a comparison of relative separations and intensities of at least two adjacent signals with pattern recognition templates, can be used as informative variables. Furthermore, this invention relates to a method and system for the on-the-fly resolution and extraction of information of signals contained in a digitized data stream involving calculation of the smoothed second

derivative of a data point from the smoothed first derivative of the

data point.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

=> d 18 kwic

SUMM

ANSWER 1 OF 1 USPATFULL L8 A method and system for automated nucleic acid sequence determination of AB a polynucleotide, wherein a nucleic acid sequencing ladder comprises signals corresponding to oligonucleotides formed from the polynucleotide, comprising the step of correlating, particularly in a trained neural network or a scatter plot, an intensity variable for each signal in the nucleic acid sequencing ladder with an informative variable for that signal, wherein the informative variable comprises information from at least two adjacent signals in the nucleic acid sequencing ladder, such that each signal in the nucleic acid sequencing ladder identified so as to determine the nucleic acid sequence corresponding to the polynucleotide. In particular, the relative separation between. SUMM . the nucleic acid sequence and a process for extracting information on-the-fly from a digitized data stream that corresponds to the nucleic acid sequencing ladder. Because the novel informative variables contain information concerning the identity of neighboring nucleotides in a polynucleotide, these variables can. . . process for resolving signals provided by this invention allows for more accurate detection of the characteristics of signals in a nucleic acid sequencing ladder, thus providing a method and system for accurately measuring these novel informative variables. Because the measurement and correlation of. SUMM . a method and system for automated nucleic acid sequence determination in which an intensity variable for each signal in the nucleic acid sequencing ladder is correlated with an informative variable for that signal such that each signal in the nucleic acid sequencing ladder is identified to determine the nucleic acid sequence corresponding to the polynucleotide. The informative variable is a variable that comprises information from at least two adjacent signals in the nucleic acid sequencing ladder, such as the relative separation between two consecutive signals, the relative intensity between two consecutive signals, or a combination. The electrophoretic separation of oligonucleotides in nucleic SUMM acid sequencing gels is primarily a function of length. However, as disclosed by this invention, the terminal nucleotides of an oligomer affect. . . mobility in a determinable fashion. Thus, the relative separation between adjacent signals, as determined for a particular signal in a nucleic acid sequencing ladder, contains information regarding the identity of terminal nucleotides, particularly the last 2-3 nucleotides, of the oligomer that corresponds to. . system for enhanced resolution of signals contained in a SUMM digitized data stream comprising successive signals corresponding to oligonucleotides in a nucleic acid sequencing ladder formed from a nucleotide. According to this method, the second derivative smoothed over m data points of the first. . . stream. Because this time-linear process provides for rapid data processing, this method is also ideally suited in contexts other than nucleic acid sequencing, such as time-of-flight mass spectrometry.

. . . for nucleic acid, particularly DNA, sequence determination of a nucleotide by correlating an intensity variable for each signal in a

```
nucleic acid sequencing ladder with an
       informative variable for that signal, wherein the informative variable
       comprises information from at least two adjacent signals in the
      nucleic acid sequencing ladder. The use of
       relative separation, relative intensity and a pattern recognition factor
       as informative variables in this method, either.
SUMM
            . a polynucleotide from a digitized data stream in which the data
       stream comprises successive signals corresponding to oligonucleotides in
       a nucleic acid sequencing ladder formed
       from a polynucleotide. Thus, it is an object of this invention to
      provide a system including means for.
       . . both streams divided by the sum of the intensities in both
DETD
       streams, can be obtained easily. For instance, in the nucleic
       acid sequencing context, the intensity of a signal
       from one data channel corresponds to the yield of a particular oligomer
       and the.
DETD
       Thus, the on-the-fly second derivative procedure does not
       destroy the fluorescence spectrochemical integrity
       that allows primary identification of the various bases. However, in
       addition to treating the fluorescence labeling of signals.
DETD
       . . . of the primary data streams provides partially deconvolved
       arrays of fluorescence and mobility data for sequences of oligomers in
       the nucleic acid sequencing context.
       Furthermore, this method is also very effective in a wide variety of
       applications beyond DNA sequencing analysis, particularly in.
DETD
       Use of Electrophoretic Separations Between Successive Oligomers as an
       Informative Variable in Nucleic Acid
       Sequencing
DETD
       The instant invention provides for an improved method and apparatus for
       nucleic acid sequencing by identifying and
       implementing new informative components of nucleic
       acid sequencing ladders in addition to the primary
       determinative variables such as lane position or fluorescence ratios.
       One such informative variable is the relative separations of successive
       oligomers in the nucleic acid sequencing
       ladders. As used herein, the term "relative separation(s)" for an
       adjacent pair of oligomers refers to either the spatial distance.
DETD
       Electrophoretic separation of oligonucleotides in nucleic
       acid sequencing gels, such as denaturing
      polyacrylamide gels, is primarily a function of length-dependent
      mobility. However, as is described in our paper,. .
DETD
      Use of Relative Oligomer Yields as an Informative Variable in
      Nucleic Acid Sequencing
            . in FIG. 5, not only does the local nucleic acid sequence affect
DETD
       the relative separations of successive oligomers in the nucleic
       acid sequencing ladders, a particular nucleotide also
       affects the yields of oligomers at neighboring positions. FIG. 5
       overlays the signal profiles from.
CLM
      What is claimed is:
       1. A method for the nucleic acid sequence determination of a
      polynucleotide, wherein a nucleic acid
       sequencing ladder comprises signals corresponding to
      oligonucleotides formed from the polynucleotide, comprising the step of
      correlating an intensity variable for each signal in the nucleic
      acid sequencing ladder with an informative variable
       for that signal, wherein the informative variable comprises information
       from at least two adjacent signals from other than a tri-nucleotide
      palindrome in the nucleic acid sequencing
       ladder, such that each signal in the nucleic acid
      sequencing ladder is identified so as to determine the nucleic
      acid sequence corresponding to the polynucleotide.
```

=>

. acid sequencer, wherein the data stream comprises successive signals corresponding to oligonucleotides from other than a tri-nucleotide palindrome in a nucleic acid sequencing ladder formed from the polynucleotide; b) locating the position of the maximum for each signal in the data stream; c). . . data stream, wherein the informative variable comprises position of maxima or intensities for at least two adjacent signals in the nucleic acid sequencing ladder; and c) correlating the informative variable for each signal with the intensity variable for each signal to determine the. . .

. comprising the steps of acquiring an intensity variable and at least one informative variable for each signal contained in a nucleic acid sequencing ladder corresponding to the polynucleotide and correlating in a trained neural network the intensity variable and the at least one. . .

. comprising the steps of acquiring an intensity variable and at least one informative variable for each signal contained in a **nucleic** acid sequencing ladder corresponding to the polynucleotide and correlating in a trained neural network the intensity variable and the at least one. . .

stream, wherein the digitized data stream comprises successive signals corresponding to oligonucleotides from other than a tri-nucleotide palindrome in a nucleic acid sequencing ladder formed from the polynucleotide, comprising: a) means for calculating an intensity variable for each signal in the digitized data. . . signal in the digitized data stream, wherein the informative variable comprises information from at least two adjacent signals in the nucleic acid sequencing

ladder; and c) means for correlating the intensity for each signal with the informative variable for each signal to identify each signal in the nucleic acid sequencing ladder so as to determine the nucleic acid sequence corresponding to the polynucleotide.

. acid sequencer, wherein the data stream comprises successive signals corresponding to oligonucleotides from other than a tri-nucleotide palindrome in a nucleic acid sequencing ladder formed from the polynucleotide; b) means for calculating the position of the maximum for each signal; c) a memory. . . signal in the digitized data stream, wherein the informative variable comprises information from at least two adjacent signals in the nucleic acid sequencing ladder; g) means for storing in the memory the informative variable for each signal; h) means for retrieving the intensity. . . correlate the intensity variable for each signal and the informative variable for each signal to identify each signal in the nucleic acid sequencing ladder so as

to determine the nucleic acid sequence corresponding to the polynucleotide.